

## BUNDESREPUBLIK DEUTSCHLAND

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02.12.1998



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DOCUMENT**  
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**Bescheinigung**

REC	10 DEC 1998
WFO	1

Die Firma Memorec Stoffel GmbH in Köln/Deutschland hat eine Patentanmeldung unter der Bezeichnung

"Neutrale Sphingomyelinase der Maus und des Menschen und diese überexprimierende Säuger- und Nicht-Säugerrzellen"

als Zusatz zur Patentanmeldung 197 34 764.9

am 15. Oktober 1997 beim Deutschen Patentamt eingereicht.

Das angeheftete Stück ist eine richtige und genaue Wiedergabe der ursprünglichen Unterlage dieser Patentanmeldung.

Die Anmeldung hat im Deutschen Patentamt vorläufig die Symbole C 12 N, C 12 Q und G 01 N der Internationalen Patentklassifikation erhalten.

München, den 22. Oktober 1998  
Der Präsident des Deutschen Patentamts  
Im Auftrag

Agurks



Aktenzeichen n: 197 58 501.9

~~HS~~  
murine nSMase

1 GTGCTGGTGG AAGCCGAGCC GGGAACAAAGG GAGGAACCTG TAGGMCGCGG  
51 TGC GGAGAAC CCACCGAAGG ACCTAAGAAT CTGGAACAGT CCACCCGAGA  
101 TTCCTTCAG GACTGCCGGC GGCCTCGCGC AGCCAGCCCCG GGATTTGCAG  
151 CCGACCTTCT TTCCGGGTGG AAGGACGGCC TTTGTCCCAG TAACGCAGGA  
201 GTAGCCCCCC ACCCCCAACC AGCTCGCGTT CCTGGGTCGG GGCAGCGCAG  
251 GACAGGGCAA TAAGCCTGTG CGCGCAATCC GCCTCGCCGC CCTTGCTCCG  
301 AAGCACTCCA GCCATGAAGC TCAACTTTTC TCTACGGCTG AGAGTTTCA  
351 ATCTCAACTG CTGGGACATC CCCTACCTGA GCAAACATAG GGCGGACCGC  
401 ATGAAGCGCT TGGGAGACTT TCTGAACCTG GAAAACCTTG ATCTGGCTCT  
451 CCTGGAGGAG GTGTGGAGTG AGCAGGACTT CCAGTACCTA AGGCAAAGGC  
501 TATCGCTCAC CTATCCAGAT GCACACTACT TCAGAAGCGG GATGATAGGC  
551 AGTGGCCTCT GTGTGTTCTC CAAACACCCA ATCCAGGAAA TCTTCCAGCA  
601 TGTCTACAGT CTGAATGGTT ACCCCTACAT GTTCCATCAT GGAGACTGGT  
651 TCTGTGGAA GTCTGTGGGG CTGCTGGTGC TCCGTCTAAG TGGACTGGTG  
701 CTCATGCCT ACCTGACTCA TCTACATGCT GAGTACAGGCC GACAGAAGGA  
751 CATCTACTTT GCACACCGTG TGGCCCAAGC TTGGGAACTG GCCCAGTCA  
801 TCCACCAACAC ATCCAAGAAT GCAGATGTGG TTCTATTGTG TGGAGACCTC  
851 AATATGCACC CCAAAGACCT GGGCTGCTGC CTGCTGAAAG AGTGGACAGG  
901 GCTCCATGAT GCTTCGTTG AGACTGAGGA CTTTAAGGGC TCTGATGATG  
951 GCTGTACCAT GGTACCCAAG AACTGCTACG TCAGCCAGCA GGACCTGGGA  
1001 CCGTTCCGT CTGGTATCCG GATTGATTAC GTGCTTTACA AGGCAGTCTC  
1051 TGAGTCCCAC GTCTGCTGTG AGACTCTGAA AACCACTACA GGCTGTGACC  
1101 CTCACAGTGA CAAGCCCTTC TCTGATCACG AGGCCCTCAT GGCTACTTTG  
1151 TATGTGAAGC ACAGCCCCCC TCAGGAAGAC CCCTGTACTG CCTGTGGCCC  
1201 ACTGGAAAGG TCCGATTTGA TCAGCGTGCT AAGGGAGGCC AGGACAGAGC  
1251 TGGGGCTAGG CATA GCTAAA GCTCGCTGGT GGGCTGCATT CTCTGGCTAT  
1301 GTGATCGTTT GGGGGCTGTC CCTTCTGGTG TTGCTGTGTG TCCTGGCTGC  
1351 AGGAGAAGAG GCCAGGGAAAG TGGCCATCAT CCTCTGCATA CCCAGTGTGG  
1401 GTCTGGTGCT GGTAGCAGGT GCAGTCTACC TCTTCCACAA GCAGGAGGCC  
1451 AAGGGCTTAT GTCGGGCCCA GGCTGAGATG CTGCACGTTT TGACAAGGGA  
1501 AACGGAGACC CAGGACCGAG GCTCAGAGCC TCACCTAGCC TACTGCTTGC  
1551 AGCAGGAGGG GGACAGAGCT TAAGAGCTTA ACAATAAAAC TTGCTTGACA  
1601 CACAAAAAAA AAAAAAAA AAAAAAAA AAA

AS  
6

human nSMase

1 GACCGCCGGG GACGAGCTTG GAGGAAAAGG AACCGGGAGC CGCCCACCCG  
51 GGGGCGCTCT CCGGACCCCC AGGGTCTAG CGCGCGGCC TTACCGAGCC  
101 TGGGCGCCCG GATTTCGGGA GCGGATCGCC TTTCCGGGTT GGCGGCCCGC  
151 CTGATTGGGA ACAGCCGGCC GGTGCGGGGG GAAACGCGGG AGTCGGGCC  
201 GACCTGAGCC ACCGCGGGCTT GGTGCCACC TGTGCGCGCC GCCTGCGAAG  
251 AAGGAACGGT CTAGGGAGAA GGCGCCGCCG GCCGCCCGC TCCCCACCGC  
301 GGCGTCGCT GGAGAGTTCG AGCCGCCTAG CGCCCGCTGGA GCTCCCCAAC  
351 CATGAAGCTC AACTTCTCCC TGCGACTGCG GATCTTCAAC CTCAACTGCT  
401 GGGGCATTCC GTACTTGAGC AAGCACCGG CCGACCGCAT GAGGCGCCGT  
451 GGAGACTTTC TGAACCAGGA GAGCTTCGAC CTGGCTTGC TGGAGGAGGT  
501 GTGGAGTGAN CAGGACTTCC AGTACCTGAG ACAGAAGCTG TCACCTACCT  
551 ACCCAGCTGC ACACCACTTC CGGAGCGGAA TCATTGGCAG TGGCCTCTGT  
601 GTCTTCTCCA AACATCCAAT CCAGGAGCTT ACCCAGCACA TCTACACTCT  
651 CAATGGCTAC CCCTACATGA TCCATCA~~tgg~~ tgactggttc agTGGGAAGG  
701 CTGTGGGCT GCTGGTGCTC CATCTAAGTG GCATGGTGCT CAACGCCAT  
751 GTGACCCATC TCCATGCCGA ATACAATCGA CAGAAGGACA TCTACCTAGC  
801 ACATCGTGTG GCCCAAGCTT GGGATTGGC CCAGTTCATC CACCACACAT  
851 CCAAGAAGGC AGACGTGGTT CTGTTGTGTG GAGACCTCAA CATGCACCCA  
901 GAAGACCTGG GCTGCTGCCT GCTGAAGGAG TGGACAGGGC TTCATGATGC  
951 CTATCTTGAA ACTCGGGACT TCAAGGGCTC TGAGGAAGGC AACACAATGG  
1001 TACCCAAGAA CTGCTACGTC AGCCAGCAGG AGCTGAAGCC ATTTCCCTTT  
1051 GGTGTCGCA TTGACTACGT GCTTTACAAG GCAGTTCTG GGTTTACAT  
1101 CTCCTGTAAG AGTTTGAAA CCACTACAGG CTTTGACCCCT CACAGTGGCA  
1151 CCCCCCTCTC TGATCATGAA GCCCTGATGG CTACTCTGTT TGTGAGGCAC  
1201 AGCCCCCCCAC AGCAGAACCC CAGCTCTACC CACGGACCAG CAGAGAGGTC  
1251 GCCGTTGATG TGTGTGCTAA AGGAGGCCG GACGGAGCTG GGTCTGGCA  
1301 TGGCTCAGGC TCGCTGGTGG GCCACCTTCG CTAGCTATGT GATTGGCCTG  
1351 GGGCTGCTTC TCTGGCACT GCTGTGTGTC CTGGCGGCTG GAGGAGGGC  
1401 CGGGGAAGCT GCCATACTGC TCTGGACCCC CAGTGTAGGG CTGGTGCTGT  
1451 GGGCAGGTGC ATTCTACCTC TTCCACGTAC AGGAGGTCAA TGGCTTATAT  
1501 AGGGCCCAGG CTGAGCTCCA GCATGTGCTA GGAAGGGCAA GGGAGGCCA  
1551 GGATCTGGC CCAGACCCCT AGCCAGCCCT ACTCCTGGGG CAGCAGGAGG  
1601 GGGACAGAAC TAAAGAACAA TAAAGCTTGG CCCTTAAAAA AAAAAAAAAA  
1651 AAAA

20  
1

mouse nSMase transl. [313 to 1572] -&gt; 1-phase Translation

DNA sequence 1632 b.p. GTGCTGGTGGAA ... AAAAAAAA linear

313 / 1 343 / 11  
ATG AAG CTC AAC TTT TCT CTA CGG CTG AGA GTT TTC AAT CTC AAC TGC TGG GAC ATC CCC  
met lys leu asn phe ser leu arg leu arg val phe asn leu asn cys trp asp ile pro  
373 / 21 403 / 31  
TAC CTG AGC AAA CAT AGG GCG GAC CGC ATG AAG CGC TTG GGA GAC TTT CTG AAC TTG GAA  
tyr leu ser lys his arg ala asp arg met lys arg leu gly asp phe leu asn leu glu  
433 / 41 463 / 51  
AAC TTT GAT CTG GCT CTC CTG GAG GAG GTG TGG AGT GAG CAG GAC TTC CAG TAC CTA AGG  
asn phe asp leu ala leu leu glu val trp ser glu gln asp phe gln tyr leu arg  
493 / 61 523 / 71  
CAA AGG CTA TCG CTC ACC TAT CCA GAT GCA CAC TAC TTC AGA AGC GGG ATG ATA GGC AGT  
gln arg leu ser leu thr tyr pro asp ala his tyr phe arg ser gly met ile gly ser  
553 / 81 583 / 91  
GGC CTC TGT GTG TTC TCC AAA CAC CCA ATC CAG GAA ATC TTC CAG CAT GTC TAC AGT CTG  
gly leu cys val phe ser lys his pro ile gln glu ile phe gln his val tyr ser leu  
613 / 101 643 / 111  
AAT GGT TAC CCC TAC ATG TTC CAT CAT GGA GAC TGG TTC TGT GGG AAG TCT GTG GGG CTG  
asn gly tyr pro tyr met phe his his gly asp trp phe cys gly lys ser val gly leu  
673 / 121 703 / 131  
CTG GTG CTC CGT CTA AGT GGA CTG GTG CTC AAT GCC TAC GTG ACT CAT CTA CAT GCT GAG  
1 leu arg leu ser gly leu val leu asn ala tyr val thr his leu his ala glu  
141 763 / 151  
CGA CAG AAG GAC ATC TAC TTT GCA CAC CGT GTG GCC CAA GCT TGG GAA CTG GCC  
tyr ser arg gln lys asp ile tyr phe ala his arg val ala gln ala trp glu leu ala  
793 / 161 823 / 171  
CAG TTC ATC CAC CAC ACA TCC AAG AAT GCA GAT GTG GTT CTA TTG TGT CGA GAC CTC AAT  
gln phe ile his his thr ser lys asn ala asp val val leu leu cys gly asp leu asn  
853 / 181 883 / 191  
ATG CAC CCC AAA GAC CTG GGC TGC TGC CTG CTG AAA GAG TGG ACA GGG CTC CAT GAT GCT  
met his pro lys asp leu gly cys cys leu leu lys glu trp thr gly leu his asp ala  
913 / 201 943 / 211  
TTC GTT GAG ACT GAG GAC TTT AAG GGC TCT GAT GTC GGC TGT ACC ATG GTA CCC AAG AAC  
phe val glu thr glu asp phe lys gly ser asp asp gly cys thr met val pro lys asn  
973 / 221 1003 / 231  
TGC TAC GTC AGC CAG CAG GAC CTG GGA CCG TTT CCG TCT GGT ATC CGG ATT GAT TAC GTG  
cys tyr val ser gln gln asp leu gly pro phe pro ser gly ile arg ile asp tyr val  
1033 / 241 1063 / 251  
CTT TAC AAG GCA GTC TCT GAG TTC CAC GTC TGC TGT GAG ACT CTG AAA ACC ACT ACA GGC  
leu tyr lys ala val ser glu phe his val cys cys glu thr leu lys thr thr gly  
1093 / 261 1123 / 271  
TGT GAC CCT CAC AGT GAC AAG CCC TTC TCT GAT CAC GAG GCC CTC ATG GCT ACT TTG TAT  
cys asp pro his ser asp lys pro phe ser asp his glu ala leu met ala thr leu tyr  
1153 / 281 1183 / 291  
CAC AGC CCC CCT CAG GAA GAC CCC TGT ACT GCC TGT GGC CCA CTG GAA AGG TCC  
his ser pro pro gln glu asp pro cys thr ala cys gly pro leu glu arg ser  
1273 / 301 1243 / 311  
GAT TTG ATC AGC GTG CTA AGG GAG GGC AGG ACA GAG CTG GGG CTA GGC ATA GCT AAA GCT  
asp leu ile ser val leu arg glu ala arg thr glu leu gly leu gly ile ala lys ala  
1273 / 321 1303 / 331  
CGC TGG TGG GCT GCA TTC TCT GGC TAT GTG ATC GTT TGG GGG CTG TCC CTT CTG GTG TTG  
arg trp trp ala ala phe ser gly tyr val ile val trp gly leu ser leu leu val leu  
1333 / 341 1363 / 351  
CTG TGT GTC CTG GCT GCA GGA GAA GAG GCC AGG GAA GTG GGC ATC ATC CTC TGC ATA CCC  
leu cys val leu ala ala gly glu glu ala arg glu val ala ile ile leu cys ile pro  
1393 / 361 1423 / 371  
AGT GTG GGT CTG GTG CTG GTA GCA GGT GCA GTC TAC CTC TTC CAC AAG CAG GAG GCC AAG  
ser val gly leu val leu val ala gly ala val tyr leu phe his lys gln glu ala lys  
1453 / 381 1483 / 391  
GGC TTA TGT CGG GCC CAG GCT GAG ATG CTG CAC GTT CTG ACA AGG GAA ACG GAG ACC CAG  
gly leu cys arg ala gln ala glu met leu his val leu thr arg glu thr glu thr gln  
1513 / 401 1543 / 411  
GAC CGA GGC TCA GAG CCT CAC CTA GCC TAC TGC TTG CAG CAG GAG GGG GAC AGA GCT TAA  
asp arg gly ser glu pro his leu ala tyr cys leu gln gln glu gly asp arg ala OCH

human nSMase transl. [352 to 1623] -&gt; 1-phase Translation

DNA sequence 1654 b.p. GACCGCCGGGGA ... AAAAAAAA linear

352 / 1 382 / 11  
ATG AAG CTC AAC TTC TCC CTG CGA CTG CGG ATC TTC AAC CTC AAC TGC TGG GGC ATT CCG  
met lys leu asn phe ser leu arg leu arg ile phe asn leu asn cys trp gly ile pro  
412 / 21 442 / 31  
TAC TTG AGC AAG CAC CGG GCC GAC CGC ATG AGG CGC CTG GGA GAC TTT CTG AAC CAG GAG  
tyr leu ser lys his arg ala asp arg met arg arg leu gly asp phe leu asn gln glu  
472 / 41 502 / 51  
AGC TTC GAC CTG GCT TTG CTG GAG GTG TGG AGT GAG CAG GAC TTC CAG TAC CTG AGA  
ser phe asp leu ala leu leu glu val trp ser glu gln asp phe gln tyr leu arg  
532 / 61 562 / 71  
CAG AAG CTG TCA CCT ACC TAC CCA GCT GCA CAC CAC TTC CGG AGC GGA ATC ATT GGC AGT  
gln lys leu ser pro thr tyr pro ala ala his his phe arg ser gly ile ile gly ser  
592 / 81 622 / 91  
GGC CTC TGT GTC TTC TCC AAA CAT CCA ATC CAG GAG CTT ACC CAG CAC ATC TAC ACT CTC  
gly leu cys val phe ser lys his pro ile gln glu leu thr gln his ile tyr thr leu  
652 / 101 682 / 111  
ATT GGC TAC CCC TAC ATG ATC CAT CAt ggt gac tgg ttc agt GGG AAG GCT GTG GGG CTG  
asn gly tyr pro tyr met ile his his gly asp trp phe ser gly lys ala val gly leu  
712 / 121 742 / 131  
CTG GTG CTC CAT CTA AGT GGC ATG GTG CTC AAC GCC TAT GTG ACC CAT CTC CAT GCC GAA  
ala leu his leu ser gly met val leu asn ala tyr val thr his leu his ala glu  
141 802 / 151  
CGA CAG AAG GAC ATC TAC CTA GCA CAT CGT GTG GCC CAA GCT TGG GAA TTG GCC  
tyr asn arg gln lys asp ile tyr leu ala his arg val ala gln ala trp glu leu ala  
832 / 161 862 / 171  
CAG TTC ATC CAC CAC ACA TCC AAG AAG GCA GAC GTG GTT CTG TTG TGT GGA GAC CTC AAC  
gln phe ile his his thr ser lys lys ala asp val val leu leu cys gly asp leu asn  
892 / 181 922 / 191  
ATG CAC CCA GAA GAC CTG GGC TGC TGC CTG CTC AAG GAG TGG ACA GGG CTT CAT GAT GCC  
met his pro glu asp leu gly cys cys leu lys glu trp thr gly leu his asp ala  
952 / 201 982 / 211  
TAT CTT GAA ACT CGG GAC TTC AAG GGC TCT GAG GAA CGC AAC ACA ATG GTA CCC AAG AAC  
tyr leu glu thr arg asp phe lys gly ser glu glu gly asn thr met val pro lys asn  
1012 / 221 1042 / 231  
TGC TAC GTC AGC CAG CAG GAG CTG AAG CCA TTT CCC TTT GGT GTC CGC ATT GAC TAC GTG  
cys tyr val ser gln gln glu leu lys pro phe pro phe gly val arg ile asp tyr val  
1072 / 241 1102 / 251  
CTT TAC AAG GCA GTT TCT GGG TTT TAC ATC TCC TGT AAG AGT TTT GAA ACC ACT ACA GGC  
leu tyr lys ala val ser gly phe tyr ile ser cys lys ser phe glu thr thr thr gly  
1132 / 261 1162 / 271  
TTT GAC CCT CAC AGT GGC ACC CCC CTC TCT GAT CAT GAA GCC CTG ATG GCT ACT CTG TTT  
phe asp pro his ser gly thr pro leu ser asp his glu ala leu met ala thr leu phe  
1192 / 281 1222 / 291  
CAC AGC CCC CCA CAG CAG AAC CCC AGC TCT ACC CAC GGA CCA GCA GAG AGG TCG  
his ser pro pro gln gln asn pro ser ser thr his gly pro ala glu arg ser  
1282 / 301 1282 / 311  
CCG TTG ATG TGT GTG CTA AAG GAG GGC TGG ACG GAG CTG CGT CTG GGC ATG GCT CAG GCT  
pro leu met cys val leu lys glu ala trp thr glu leu gly leu gly met ala gln ala  
1312 / 321 1342 / 331  
CCG TGG TGG GCC ACC TTC GCT AGC TAT GTG ATT CGC CTG GGG CTG CTT CTC CTG GCA CTG  
arg trp trp ala thr phe ala ser tyr val ile gly leu gly leu leu leu ala leu  
1372 / 341 1402 / 351  
CTG TGT GTC CTG GCG GCT CGA GGA GGG GCC GGG GAA GCT GCC ATA CTG CTC TGG ACC CCC  
leu cys val leu ala ala gly gly ala gly glu ala ala ile leu leu trp thr pro  
1432 / 361 1462 / 371  
AGT GTA GGG CTG GTG CTG TGG GCA GGT GCA TTC TAC CTC TTC CAC GTA CAG GAG GTC AAT  
ser val gly leu val leu trp ala gly ala phe tyr leu phe his val gln glu val asn  
1492 / 381 1522 / 391  
GGC TTA TAT AGG GCC CAG GCT GAG CTC CAG CAT GTG CTA GGA AGG GCA AGG GAG GGC CAG  
gly leu tyr arg ala gln ala glu leu gln his val leu gly arg ala arg glu ala gln  
1552 / 401 1582 / 411  
GAT CTG GGC CCA GAG CCT CAG CCA GCC CTA CTC CTG GGG CAG CAG GAG GGG GAC AGA ACT  
asp leu gly pro glu pro gln pro ala leu leu gly gln gln glu gly asp arg thr  
1612 / 421  
AAA GAA CAA TAA  
lys glu gln OCH

4  
9

801 ACATCGTGTG GCCCAAGCTT GGGATTGGC CCAGTTICATC CACCACACAT  
851 CCAAGAAGGC AGACGTGGTT CTGTTGTGTG GAGACCTCAA CATGCACCCA  
901 GAAGACCTGG GCTGCTGCCT GCTGAAGGAG TGGACAGGGC TTCATGATGC  
951 CTATCTTGAA ACTCGGGACT TCAAGGGCTC TGAGGAAGGC AACACAATGG  
1001 TACCCAAAGAA CTGCTACGTC AGCCAGCAGG AGCTGAAGCC ATTTCCCTTT  
1051 GGTGTCCGCA TTGACTACGT GCTTTACAAG GCAGTTTCTG GGTTTTACAT  
1101 CTCCCTGTAAG AGTTTTGAAA CCACTACAGG CTTTGACCCCT CACAGTGGCA  
1151 CCCCCCTCTC TGATCATGAA GCCCTGATGG CTACTCTGTT TGTGAGGCAC  
1201 AGCCCCCCCAC AGCAGAACCC CAGCTCTACC CACGGACCAG CAGAGAGGTC  
1251 GCCGTTGATG TGTGTGCTAA AGGAGGCCTG GACGGAGCTG GGTCTGGCA  
1301 TGGCTCAGGC TCGCTGGTGG GCCACCTTCG CTAGCTATGT GATTGGCCTG  
1351 GGGCTGCTTC TCCTGGCACT GCTGTGTGTC CTGGCGGCTG GAGGAGGGC  
1401 CGGGGAAGCT GCCATACTGC TCTGGACCCC CAGTGTAGGG CTGGTGCTGT  
1451 GGGCAGGTGC ATTCTACCTC TTCCACGTAC AGGAGGTCAA TGGCTTATAT  
1501 AGGGCCCCAGG CTGAGCTCCA GCATGTGCTA GGAAGGGCAA GGGAGGCCA  
1551 GGATCTGGGC CCAGAGCCTC AGCCAGCCCT ACTCCTGGGG CAGCAGGAGG  
1601 GGGACAGAAC TAAAGAACAA TAAAGCTTGG CCCTTTAAAA AAAAAAAA  
1651 AAAA

underlined is the coding sequence

25  
10

## mouse nSMase

1 GTGCTGGTGG AAGCCGAGCC GGGAAACAAGG GAGGAACCTG TAGGMCGCG  
 51 TCGGGAGAAC CCACCGAAGG ACCTAAGAAT CTGGAACAGT CCACCCGAGA  
 101 TTCCCTTCAG GACTGCCGC GCCCTCGCGC AGCCAGCCCG GGATTTCAG  
 151 CCGACCTTCT TTCCGGGTGG AAGGACGGCC TTGTCAGGAGA TAACCGAGGA  
 201 GTAGCCCCCC ACCCCCAACC AGCTCGCGTT CCTGGGTGCG GGCAGCGAG  
 251 GACAGGGCAA TAAGCCTGTG CGCGCAATCC GCCTCGCCGC CCTTGCTCCG  
 301 AAGCACTCCA GCCATGAAGC TCAACTTTTC TCTACGGCTG AGAGTTTCA  
 351 ATCTCAACTG CTGGGACATC CCCTACCTGA GCAAACATAG GGCAGGACCGC  
 401 ATGAAGCGCT TGGGAGACTT TCTGAACCTTG GAAAACCTTG ATCTGGCTCT  
 451 CCTGGAGGAG GTGTGGAGTG AGCAGGACTT CCAGTACCTA AGGCAAAGGC  
 501 TATCGCTCAC CTATCCAGAT GCACACTACT TCAGAAGCGG GATGATAGGC  
 551 AGTGGCCTCT GTGTGTTCTC CAAACACCCA ATCCAGGAAA TCTTCCAGCA  
 601 TGTCTACAGT CTGAATGGTT ACCCCTACAT GTTCCATCAT GGAGACTGGT  
 651 TCTGTGGAA GTCTGTGGGG CTGCTGGTGC TCCGTCTAAG TGGACTGGTG  
 701 CTCAAATGCCT ACGTGACTCA TCTACATGCT GAGTACAGCC GACAGAAGGA  
 751 CATCTACTTT GCACACCGTG TGGCCCAAGC TTGGGAACCTG GCCCAGTTCA  
 801 TCCACCCACAC ATCCAAGAAT GCAGATGTGG TTCTATTGTG TGGAGACCTC  
 851 AATATGCACC CCAAAGACCT GGGCTGCTGC CTGCTGAAAG AGTGGACAGG  
 901 GCTCCATGAT GCTTCGTTG AGACTGAGGA CTTTAAGGGC TCTGATGATG  
 951 GCTGTACCAT GGTACCCAAG AACTGCTACG TCAGCCAGCA GGACCTGGGA  
 1001 CCGTTCCGT CTGGTATCCG GATTGATTAC GTGCTTTACA AGGCAGTCTC  
 1051 TGAGTTCCAC GTCTGCTGTG AGACTCTGAA ACCACTACA GGCTGTGACC  
 1101 CTCACAGTGA CAAGCCCTTC TCTGATCACG AGGCCCTCAT GGCTACTTTG  
 1151 TATGTGAAGC ACAGCCCCCCC TCAGGAAGAC CCCTGTACTG CCTGTGGCCC  
 1201 ACTGGAAAGG TCCGATTGTA TCAGCGTGCT AAGGGAGGCC AGGACAGAGC  
 1251 TGGGGCTAGG CATAGCTAAA GCTCGCTGGT GGGCTGCATT CTCTGGCTAT  
 1301 GTGATCGTTT GGGGGCTGTC CCTTCTGGTG TTGCTGTG TGCTGGCTGC  
 1351 AGGAGAAAGAG GCCAGGGAAG TGGCCATCAT CCTCTGCATA CCCAGTGTGG  
 1401 GTCTGGTGCT GGTAGCAGGT GCAGTCTACC TCTCCACAA GCAGGAGGCC  
 1451 AAGGGCTTAT GTCGGGCCCA GGCTGAGATG CTGCACTGTT TGACAAGGGG  
 1501 AACGGAGACC CAGGACCGAG GCTCAGAGCC TCACCTAGCC TACTGCTTGC  
 1551 AGCAGGAGGG GGACAGAGCT TAAGAGCTTA ACAATAAAAC TTGCTTGACA  
 1601 CACAAAAAAA AAAAAAAA AAAAAAAA AAA

## human nSMase

1 GACCGCCGGG GACGAGCTTG GAGGAAAAGG AACCGGGAGC CGCCCACCCG  
 51 GGGGCGCTCT CGGGACCCCC AGGGTCCTAG CGCGCGGCC TTACCGAGCC  
 101 TGGGCGCCCG GATTTGGGA GCGGATCGCC TTTCCGGTTT GGCAGGCCGC  
 151 CTGATTGGGA ACAGCCGGCC GGTTGCCGGG GGAACGCCGG AGTCGGGCC  
 201 GACCTGAGCC ACAGCGGGCTT GGTGCCCACC TGTGCGCGCC GCCTGCGAAG  
 251 AAGGAACGGT CTAGGGAGAA GGCGCCGCCG GCCGCCCCCG TCCCCACCGC  
 301 GGCCGTCGCT GGAGAGTTCG AGCCGCCTAG CGCCCTGGA GCTCCCCAAC  
 351 CATGAAGCTC AACATCTCCC TGCGACTGCG GATCTTCAAC CTCAACTGCT  
 401 GGGGCATTCC GTACTTGAGC AACGACCGGG CGCACCGCAT GAGGCGCTG  
 451 GGAGACTTTC TGAACCAGGA GAGCTTCGAC CTGGCTTTGC TGGAGGAGGT  
 501 GTGGAGTGAG CAGGACTTCC AGTACCTGAG ACAGAAGCTG TCACCTACCT  
 551 ACCCAGCTGC ACACCACTTC CGGAGCGGAA TCATTGGCAG TGGCCCTCTGT  
 601 GTCTTCTCCA AACATCCAAT CCAGGAGCTT ACCCAGCACA TCTACACTCT  
 651 CAATGGCTAC CCCTACATGA TCCATCATGA tgactaattc agTGGGAAGG  
 701 CTGTGGGCT GCTGGTGCTC CATCTAAGTG GCATGGTGCT CAACGCCTAT  
 751 GTGACCCATC TCCATGCCGA ATACAATCGA CAGAAGGACA TCTACCTAGC

